Amendments to the Claims

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

- 1.-25. (Canceled)
- 26. (Withdrawn) A method for producing a chimeric non-human animal comprising a modified foreign chromosome(s) or a fragment(s) thereof, which comprises the steps of:
- (a) preparing a microcell comprising a foreign chromosome(s) or a fragment(s) thereof, and transferring said foreign chromosome(s) or a fragment(s) thereof into a cell with high homologous recombination efficiency through its fusion with said microcell;
- (b) in said cell with high homologous recombination efficiency, inserting a vector by homologous recombination into a desired site of said foreign chromosome(s) or a fragment(s) thereof, and/or a desired site of a chromosome(s) derived from said cell with high homologous recombination efficiency, thereby marking said desired site;
- (c) in said cell with high homologous recombination efficiency, causing deletion and/or translocation to occur at the marked site of said foreign chromosome(s) or a fragment(s) thereof; and
- (d) preparing a microcell comprising said foreign chromosome(s) or a fragment(s) thereof in which deletion or translocation has occurred, and transferring said foreign chromosome(s) or a fragment(s) thereof into a pluripotent non-human animal cell through its fusion with said microcell.
- 27. (Withdrawn) The method of claim 26, wherein a plurality of said cells with high homologous recombination efficiency are subjected to whole cell fusion after steps (a) and (b) and are subjected to step (c).
- 28. (Withdrawn) The method of claim 27, wherein a plurality of said cells with high homologous recombination efficiency each comprise a distinct foreign chromosome(s) or a fragment(s) thereof.
- 29. (Withdrawn) The method of claim 26, wherein said targeting vector comprises a telomere sequence which is introduced into a desired site by insertion of the targeting vector.

- 30. (Withdrawn) The method of claim 29, wherein said deletion occurs at a site where said telomere sequence has been introduced.
- 31. (Withdrawn) The method of claim 26, wherein said targeting vector comprises a recognition sequence for a site-directed recombination enzyme, and said recognition sequence is introduced into a desired site by insertion of the targeting vector.
- 32. (Withdrawn) The method of claim 31, wherein a vector, which is capable of expressing a site-directed recombination enzyme, is introduced into said cell with high homologous recombination efficiency simultaneously with or after insertion of said targeting vector comprising said recognition sequence for a site-directed recombination enzyme, so that an activity of said site-directed recombination enzyme is expressed, resulting in deletion and/or translocation of said foreign chromosome(s) or a fragment(s) thereof at a site into which said recognition sequence is introduced.
- 33. (Withdrawn) The method of claim 32, wherein said translocation occurs between a plurality of foreign chromosomes or fragments thereof.
- 34. (Withdrawn) The method of claim 32, wherein said translocation occurs between said foreign chromosome(s) or a fragment(s) thereof and said chromosome(s) derived from said cell with high homologous recombination efficiency.
- 35. (Withdrawn) The method of claim 31, wherein said site-directed recombination enzyme is a Cre enzyme.
- 36. (Withdrawn) The method of claim 31, wherein said recognition sequence for sitedirected recombination enzyme is a LoxP sequence.
- 37. (Withdrawn) The method of claim 26, wherein said cell with high homologous recombination efficiency is an embryonic stem cell (or ES cell).
- 38. (Withdrawn) The method of claim 26, wherein said cell with high homologous recombination efficiency is a chicken DT-40 cell.
- 39. (Withdrawn) The method of claim 26, which further comprises a step of screening cells comprising a foreign chromosome(s) or a fragment(s) thereof in which deletion and/or translocation has occurred.

- 40. (Withdrawn) The method of claim 39, wherein said screening is based on expression of a marker gene.
- 41. (Withdrawn) The method of claim 40, wherein said marker gene is a drug-resistance gene.
- 42. (Withdrawn) The method of claim 40, the marker gene is a green fluorescent proteinencoding gene derived from the jellyfish Aequorea victoria or a modified gene thereof.
- 43. (Withdrawn) The method of claim 26, wherein in the step (d), a microcell is produced from said cell with high homologous recombination efficiency; said foreign chromosome(s) or a fragment(s) thereof, in which deletion and/or translocation has occurred is transferred into a CHO cell through its fusion with said microcell; a microcell is produced from the CHO cell; and then said foreign chromosome(s) or a fragment(s) thereof in which deletion and/or translocation has occurred is transferred into a pluripotent cell through its fusion with said microcell.
- 44. (Withdrawn) The method of claim 26, said pluripotent cell is an embryonic stem cell (or ES cell).
- 45. (Withdrawn) The method of claim 26, said foreign chromosome(s) or a fragment(s) thereof is derived from a human.
- 46. (Withdrawn) A method for producing a non-human animal comprising a modified foreign chromosome(s) or a fragment(s) thereof, which comprises the steps of:
- (a) preparing a microcell comprising a foreign chromosome(s) or a fragment(s) thereof, and transferring said foreign chromosome(s) or a fragment(s) thereof into a cell with high homologous recombination efficiency thorugh its fusion with said microcell;
- (b) in said cell with high homologous recombination efficiency, inserting a vector by homologous recombination into a desired site of said foreign chromosome(s) or a fragment(s) thereof, and/or a desired site of a chromosome(s) derived from said cell with high homologous recombination efficiency, thereby marking said desired site;
- (c) in said cell with high homologous recombination efficiency, causing deletion and/or translocation to occur at the marked site of said foreign chromosome(s) or a fragment(s) thereof;

- (d) preparing a microcell comprising said foreign chromosome(s) or a fragment(s) thereof, in which deletion and/or translocation has occurred, and transferring said foreign chromosome(s) or a fragment(s) thereof into a cell derived from a non-human animal through its fusion with said microcell; and
- (e) transplanting the nucleus of said cell derived from the non-human animal into an enucleated unfertilized egg derived from a homologous non-human animal of the same species.
- 47. (Withdrawn) The method of claim 46, wherein a plurality of said cells with high homologous recombination efficiency are subjected to whole cell fusion after steps (a) and (b) and are subjected to the step (c).
- 48. (Withdrawn) The method of claim 47, wherein a plurality of said cells with high homologous recombination efficiency comprise a distinct foreign chromosome(s) or a fragment(s) thereof.
- 49. (Withdrawn) The method of claim 46, wherein said targeting vector comprises a telomere sequence, which is introduced into a desired site by insertion of the targeting vector.
- 50. (Withdrawn) The method of claim 49, wherein said deletion occurs at a site into which a telomere sequence has been introduced.
- 51. (Withdrawn) The method of claim 46, wherein said targeting vector comprises a recognition sequence for site-directed recombination enzyme, and said recognition sequence is introduced into a desired site by insertion of the targeting vector.
- 52. (Withdrawn) The method of claim 51, wherein a vector, which is capable of expressing a site-directed recombination enzyme, is introduced into said cell with high homologous recombination efficiency simultaneously with or after insertion of said targeting vector comprising said recognition sequence for a site-directed recombination enzyme, so that an activity of said site-directed recombination enzyme is expressed, resulting in deletion and/or a translocation of said foreign chromosome(s) or fragment(s) thereof at a site into which said recognition sequence is introduced.

- 53. (Withdrawn) The method of claim 52, wherein said translocation occurs between a plularity of foreign chromosomes or fragments thereof.
- 54. (Withdrawn) The method of claim 52, wherein said translocation occurs between said foreign chromosome(s) or a fragment(s) thereof and said chromosome derived from a cell with high homologous recombination efficiency.
- 55. (Withdrawn) The method of claim 51, wherein said site-directed recombination enzyme is a Cre enzyme.
- 56. (Withdrawn) The method of claim 51, wherein said recognition sequence for a sitedirected recombination enzyme is a LoxP sequence.
- 57. (Withdrawn) The method of claim 46, wherein said cell with high homologous recombination efficiency is an embryonic stem cell (or ES cell).
- 58. (Withdrawn) The method of claim 46, wherein said cell with high homologous recombination efficiency is a chicken DT-40 cell.
- 59. (Withdrawn) The method of claim 46, which further comprises a step of screening cells containing a foreign chromosome(s) or a fragment(s) thereof in which deletion and/or translocation has occurred.
- 60. (Withdrawn) The method of claim 59, wherein said screening is based on expression of a marker gene.
- 61. (Withdrawn) The method of claim 60, wherein said marker gene is a drug-resistant gene.
- 62. (Withdrawn) The method of claim 60, wherein said marker gene is a green fluorescent protein-encoding gene derived from the jellyfish Aequorea victoria or a modified gene thereof.
- 63. (Withdrawn) The method of claim 46, wherein, in the step (d), a microcell is produced from said cell with high homologous recombination efficiency; said foreign chromosome(s) or fragment(s) thereof, in which deletion and/or translocation have/has occurred, is/are transferred

into a CHO cell through its fusion with the microcell; a microcell is produced from the CHO cell; and then said foreign chromosome(s) or a fragment(s) thereof, in which deletion and/or translocation has occurred, is transferred into a cell derived from a non-human animal through its fusion with the microcell.

- 64. (Withdrawn) The method of claim 46, said cell derived from a non-human animal is a culture cell derived from an embryo or a blastocyst.
- 65. (Withdrawn) The method of claim 46, said cell derived from a non-human animal is a culture cell derived from a fetus or an adult.
- 66. (Withdrawn) The method of claim 46, said cell derived from a non-human animal is a fibroblast cell derived from fetus.
- 67. (Withdrawn) The method of claim 46, said foreign chromosome(s) or a fragment(s) thereof is derived from a human.
- 68. (Withdrawn) A non-human animal, which retains a chromosomal fragment(s) obtained by deletion of a foreign chromosome(s) or a fragment(s) thereof.
- 69. (Withdrawn) The non-human animal of claim 68, wherein said chromosomal fragment(s) comprises:
 - (i) a marker gene and a telomere sequence, and/or
 - (ii) a recognition sequence for a site-directed recombination enzyme.
- 70. (Withdrawn) A non-human animal, comprising a recombinant foreign chromosome(s) obtained by translocation between a plurality of foreign chromosomes or fragments thereof.
- 71. (Withdrawn) The non-human animal of claim 70, wherein said recombinant chromosomal fragment(s) comprises:
 - (i) a marker gene and a telomere sequence; and/or
 - (ii) a recognition sequence for a site-directed recombination enzyme.

- 72. (Withdrawn) The non-human animal of claim 70, wherein said recombinant foreign chromosome(s) is independently maintained in the nucleus of the non-human animal cell.
- 73. (Withdrawn) The non-human animal of claim 70, wherein said recombinant foreign chromosome(s) is derived from a human.
- 74. (Withdrawn) The non-human animal of claim 70, wherein the recombinant foreign chromosome(s) is derived from human chromosomes #14 and #2.
- 75. (Withdrawn) The non-human animal of claim 70, wherein said recombinant foreign chromosome(s) is derived from human chromosomes #14 and #22
- 76. (Withdrawn) The non-human animal of claim 70, wherein said recombinant foreign chromosome(s) comprises genes for a heavy-chain and a light-chain of a human antibody.
- 77. (Withdrawn) The non-human animal of claim 70, wherein said recombinant foreign chromosome(s) comprises genes for a heavy-chain and a light-chain * gene of a human antibody.
 - 78. (Withdrawn) The non-human animal of claim 70, which is a mouse.
 - 79. (Withdrawn) The non-human animal of claim 70, which is an ungulata.
 - 80. (Withdrawn) The non-human animal of claim 70, which is a bovine.
 - 81. (Withdrawn) The non-human animal of claim 70, which is an ovine.
 - 82. (Withdrawn) The non-human animal of claim 70, which is an avian.
 - 83. (Withdrawn) The non-human animal of claim 70, which is a chicken.
 - 84. (Canceled)
- 85. (Withdrawn) A method for modifying a foreign chromosome(s) or a fragment(s) thereof in a cell, which comprises the steps of:
- (a) preparing a microcell containing a foreign chromosome(s) or a fragment(s) thereof, and transferring said foreign chromosome(s) or a fragment(s) thereof into a cell with high homologous recombination efficiency through its fusion with the microcell;

- (b) in said cell with high homologous recombination efficiency, inserting a targeting vector by homologous recombination into a desired site of said foreign chromosome(s) or a fragment(s) thereof and/or a desired site of a chromosome(s) derived from said cell with high homologous recombination efficiency, thereby marking said desired site; and
- (c) in said cell with high homologous recombination efficiency, causing deletion or translocation to occur at the marked site of said foreign chromosome(s) or a fragment(s) thereof.
 - 86.-92. (Canceled)
 - 93. (Currently amended) A recombinant chromosome comprising:
 - (i) the centromere of human chromosome #14;
 - (ii) two telomere sequences;
 - (iii) at least one recognition sequence for a site-directed recombination enzyme;
- (iv) at least two chromosome fragments that are <u>had</u> not <u>been</u> adjacently located in a natural chromosome or in a naturally occurring chromosome fragment; and
 - (iv) a marker gene,

wherein said recognition sequence for a site-directed recombination enzyme is located between said two chromosome fragments.

- 94. (Previously presented) The recombinant chromosome of claim 93, wherein (A) said centromere is contained within a human chromosome #14 fragment and (B) said recombinant chromosome comprises at least one chromosome fragment that is not naturally located adjacent to said human chromosome #14 fragment.
- 95. (Previously presented) The recombinant chromosome of claim 94, wherein said human chromosome #14 fragment is a centromere-comprising portion of the chromosome fragment denoted as SC20.
- 96. (Previously presented) The recombinant chromosome of claim 93, wherein one chromosome fragment is a fragment of human chromosome #2.

- 97. (Previously presented) The recombinant chromosome of claim 93, wherein one chromosome fragment is a fragment of human chromosome #22.
- 98. (Previously presented) The recombinant chromosome of claim 93, comprising a human chromosome #14 fragment and a human chromosome #2 fragment.
- 99. (Currently amended) The recombinant chromosome of claim 98, wherein said chromosome #14 fragment comprises fragments comprise a human antibody heavy-chain gene locus and said chromosome #2 fragment comprises a human antibody light-chain kappa gene locus.
- 100. (Currently amended) The recombinant chromosome of claim 98, wherein said chromosome #14 fragment comprises fragments comprise the entire region of the human antibody heavy-chain gene locus and said chromosome #2 fragment comprises the entire region of the human antibody light-chain kappa gene locus.
- 101. (Previously presented) The recombinant chromosome of claim 93, comprising a human chromosome #14 fragment and a human chromosome #22 fragment.
- 102. (Currently amended) The recombinant chromosome of claim 101, wherein said chromosome #14 fragment comprises fragments comprise a human antibody heavy-chain gene locus and said chromosome #22 fragment comprises a human antibody light-chain lambda gene locus.
- 103. (Currently amended) The recombinant chromosome of claim 101, wherein said chromosome #14 fragment comprises fragments comprise the entire region of the human antibody heavy-chain gene locus and said chromosome #22 fragment comprises the entire region of the human antibody light-chain lambda gene locus.
- 104. (Previously presented) The recombinant chromosome of claim 93, which is generated by chromosome recombination between the chromosome fragment denoted as SC20 and another chromosome fragment.
- 105. (Previously presented) The recombinant chromosome of claim 104, wherein said recombinant chromosome comprises the entire region of the human antibody heavy chain gene locus.

- 106. (Previously presented) The recombinant chromosome of claim 104, which is generated by chromosome recombination between the chromosome fragment denoted as SC20 and a fragment of a chromosome other than the human chromosome #14.
- 107. (Previously presented) The recombinant chromosome of claim 106, wherein the fragment of a chromosome other than the human chromosome #14 is a fragment of a human chromosome #2, which comprises a human antibody light-chain kappa gene locus.
- 108. (Previously presented) The recombinant chromosome of claim 106, wherein the fragment of a chromosome other than the human chromosome #14 is a fragment of human chromosome #22, which comprises a human antibody light-chain lambda gene locus.
- 109. (Previously presented) The recombinant chromosome of claim 93, which comprises both a human antibody heavy-chain gene locus and a human antibody light-chain gene locus.
- 110. (Previously presented) The recombinant chromosome of claim 93, which comprises both the entire region of the human antibody heavy-chain gene locus and the entire region of the human antibody light-chain gene locus.
- 111. (Previously presented) The recombinant chromosome of claim 93, wherein said recognition sequence is the loxP sequence and said site-directed recombination enzyme is the Cre recombinase.
- 112. (Previously presented) The recombinant chromosome of claim 93, wherein said recognition sequence is the FRP sequence and said site-directed recombination enzyme is the FLP recombinase.
 - 113. (Currently amended) A recombinant chromosome, which comprises:
 - (i) the centromere sequence of a human chromosome #21;
 - (ii) two telomere sequences;
 - (iii) at least one recognition sequence for site-directed recombination enzyme;
- (iv) at least two chromosome fragments that are <u>had</u> not <u>been</u> adjacently located in a natural chromosome <u>or in a naturally occurring chromosome fragment</u>; and

(iv) a marker gene,

wherein said recognition sequence for a site-directed recombination enzyme is located between said two chromosome fragments.

- 114. (Previously presented) The recombinant chromosome of claim 113, wherein (A) said centromere is contained within a human chromosome #21 fragment and (B) said recombinant chromosome comprises at least one chromosome fragment that is not naturally located adjacent to said human chromosome #21 fragment.
- 115. (Previously presented) The recombinant chromosome of claim 113, wherein said recognition sequence is the loxP sequence and said site-directed recombination enzyme is the Cre recombinase.
- 116. (Previously presented) The recombinant chromosome of claim 113, wherein said recognition sequence is the FRP sequence and said site-directed recombination enzyme is the FLP recombinase.
- 117. (Currently amended) A method for producing a recombinant chromosome, comprising:
- (a) preparing a first <u>isolated</u> cell comprising a fragment of human chromosome #14 that has a centromere and a recognition sequence for a site-directed recombination enzyme positioned at desired site within <u>in</u> said fragment;
- (b) preparing a second <u>isolated</u> cell comprising a second chromosome fragment, which comprises a recognition sequence for a site-directed recombination enzyme positioned at desired site in said second chromosome fragment;
 - (c) fusing said first cell with said second cell to produce a hybrid cell; and
 - (d) expressing a site-directed recombination enzyme in said hybrid cell,

wherein said enzyme causes site-directed recombination between said fragment of human chromosome #14 and said second chromosome fragment to generate a recombinant chromosome,

wherein said recombinant chromosome comprises the centromere of human chromosome #14 and a portion of the second chromosome fragment.

- 118. (Previously presented) The method of claim 117, wherein said recombinant chromosome is transferred from said hybrid cell into a new cell type via microcell fusion.
- 119. (Previously presented) The method of claim 118, wherein said new cell type is a CHO cell.
- 120. (Previously presented) The method of claim 117, wherein said first cell and said second cell are chicken DT-40 cells.
- 121. (Previously presented) The method of claim 117, wherein said site-directed recombination is detected by the expression of a reporter gene.
- 122. (Previously presented) The method of claim 121, wherein said reporter gene is a green fluorescent protein gene or functional variant thereof.
- 123. (Previously presented) The method of claim 117, wherein said recognition sequence in said human chromosome #14 fragment and said recognition sequence in said second chromosome fragment are loxP sequences, and said site-directed recombination enzyme is the Cre recombinase.
- 124. (Previously presented) The method of claim 117, wherein said recognition sequence in said human chromosome #14 fragment and said recognition sequence in said second chromosome fragment is the FRP sequence and said site-directed recombination enzyme is the FLP recombinase.
- 125. (Previously presented) The method of claim 117, said human chromosome #14 fragment is the chromosome fragment denoted as SC20.
- 126. (Previously presented) The method of claim 117, said second chromosome fragment is a fragment of either human chromosome #2 or human chromosome #22, comprising a human antibody light chain gene locus.
- 127. (Currently amended) A method for producing a recombinant chromosome, comprising:

- (a) preparing a first <u>isolated</u> cell comprising a fragment of human chromosome #21 that has a centromere and a recognition sequence for a site-directed recombination enzyme positioned at desired site within <u>in</u> said fragment;
- (b) preparing a second <u>isolated</u> cell comprising a second chromosome fragment, which comprises a recognition sequence for a site-directed recombination enzyme positioned at desired site in said second chromosome fragment;
 - (c) fusing said first cell with said second cell to produce a hybrid cell; and
 - (d) expressing a site-directed recombination enzyme in said hybrid cell,

wherein said enzyme causes site-directed recombination between said fragment of human chromosome #21 and said second chromosome fragment to generate a recombinant chromosome, wherein said recombinant chromosome comprises the centromere of human chromosome #21 and a portion of the second chromosome fragment.

- 128. (Previously presented) The method of claim 127, wherein said recombinant chromosome is transferred from said hybrid cell into a new cell type via microcell fusion.
- 129. (Previously presented) The method of claim 128, wherein said second cell is a CHO cell.
- 130. (Previously presented) The method of claim 127, wherein said first cell and said second cell are chicken DT-40 cells.
- 131. (Previously presented) The method of claim 127, wherein said site-directed recombination is detected by the expression of a reporter gene.
- 132. (Previously presented) The method of claim 131, wherein said reporter gene is a green fluorescent protein gene or functional variant thereof.
- 133. (Previously presented) The method of claim 127, wherein said recognition sequence in said human chromosome #21 fragment and said recognition sequence in said second chromosome fragment are loxP sequences, and said site-directed recombination enzyme is the Cre recombinase.

- 134. (Previously presented) The method of claim 127, wherein said recognition sequence in said human chromosome #21 fragment and said recognition sequence in said second chromosome fragment is the FRP sequence and said site-directed recombination enzyme is the FLP recombinase.
- 135. (Previously presented) A cell comprising the recombinant chromosome of any one of claims 93 or 113.
- 136. (Previously presented) The method of claim 117, wherein said recognition sequence for a site-directed recombination enzyme is positioned at said desired site in said human chromosome #14 fragment and said second chromosome fragment by a targeting vector.
- 137. (Previously presented) The method of claim 127, wherein said recognition sequence for a site-directed recombination enzyme is positioned at said desired site in said human chromosome #21 fragment and said second chromosome fragment by a targeting vector.
- 138. (Previously presented) The method of claim 122, wherein said green fluorescent protein gene or functional variant thereof, is obtained from the jellyfish *Aequorea victoria*.
- 139. (Previously presented) The method of claim 132, wherein said green fluorescent protein gene or functional variant thereof, is obtained from the jellyfish *Aequorea victoria*.
- 140. (New) A chromosome vector comprising: (i) a chromosome fragment comprising the centromere of human chromosome #21; (ii) two telomere sequences; (iii) at least one recognition sequence for a site-directed recombination enzyme; and (iv) a marker gene.
- 141. (New) The chromosome vector of claim 140, wherein said chromosome fragment is a fragment of human chromosome #21.
- 142. (New) The chromosome vector of claim 140, wherein said recognition sequence is the loxP sequence and said site-directed recombination enzyme is the Cre recombinase.
- 143. (New) The chromosome vector of claim 140, wherein said recognition sequence is the FRP sequence and said site-directed recombination enzyme is the FLP recombinase.